			0	7 - 0	3.00			
UTILITY PATENT APPLICATION					ATTORNEY DOCKET 81326D-			
TRANSMITTAL UNDER 37 CFR 1.53(b)								
Commissioner for Patents			Express Mail Label No.					
Box Patent Applica	ation							
Washington, D.C. 20231				EL 485197960US				
n							ဥ	
ETHOD OF CURING A	FUSE	R MEMB	ER	Date:	Jane	S0,2000	<u>م</u>	
VERCOAT AT LOW TH					7		900	
· Encount in a a a a			i i		O		50	
irst Named Inventor (or A	onlica	tion Identi	ifier):				oss4 U.S. PTO	
Bot I tallion III tollion (or a	-F						- S	
ann H. Chen, et al								
unii 11. Onon, ot ui			1					
nclosed are:								
X Specification				6.	X Assi	gnment of the invention to		
					Nex	Press Solutions LLC		
Sheet(s) of drawing	(s)			7.	Cert	ified copy of a priority		
Information Disclo	nea Stat	ement I Inde	er 37 CFR	8.	docu	ment ociate Power of Attorney		
1.97.	surc Stat	cincii Oliu	21 57 CIR	٠.		Jeante I o West of Tittlesiney		
Combined Declaration fo	. Dotont	Annlication	and Dower of	Attorr	ev.			
4a. X New	ratent	Аррисацон	and I ower or	Auon	cy.			
	prior an	nlication (3	7 CFR 1 63(d)	(for c	ontinuation/di	visional with Box 11 comp	leted)	
40. Copy nom a	prior ap	phoanon (5	, CI K 1.05(a)	(101 0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	The last water and the comp	,	
. Incorporation by R	eference	(useable if	Box 4b is	9.	Del	etion of Inventor(s).		
hecked) The entire disclosure				Signed statement attached deleting inventor(s) name				
hich a copy of the oath or dec				in	the prior appl	ication, see 37 CFR 1.63(d)	(2) and	
considered as being part of the	e disclo	sure of the a	ccompanying	1.3	3(b).			
pplication and is hereby incorp								
				-ident	fied application	on, amend the specification	at Page	
after the title, by insCROSS REFEREN				N				
CRUSS REFEREN	made t	o and priori	by claimed from	m ILS.	Provisional A	application Serial No.,		
filed, entitled.	, , , , , , , , , , , , , , , , , , , ,	o and priors	.,			,		
a CONTINUING APPLICA	TION,	check appro	priate box an	d suppl	y the requisit	e information:		
1. Continuation	Divisio	nal	Continuation-	in-part	(CIP) of	prior application No: ,		
2. X Please address all wr	itten cor	nmunication	ns to Lawrence	e P. Ke	ssler.			
NexPress Solutions								
Please Direct all tele	phone c	alls to Law	rence P. Kes	ssler (	716) 726-9	178.		
he filing fee has been calculat FOR:		FILED	NO. EXTRA	ίT.	RATE	FEE		
BASIC FEE		TILLE	110. 22111	-	10112	\$ 690		
OTAL CLAIMS	20	- 20 =	0		x 18 =	\$ 0		
NDEPENDENT CLAIMS 1 - 3 = 0				1.	x 78 =	\$ 0		
MULTIPLE DEPENDEN	T CLAI	M PRESEN	TED		+ 260	\$0		
				L	TOTAL	\$ 690		
	77 1 1	C	N	4 NT.	05 0225 1-4-	e amount of \$ 690.		
Please charge my Eastma			Deposit Accou ppy of this she			e amount of \$ 090.		
	A	иприсите с	ANS SILL LIES SILE	cet is e	it.ioscu			

X The Commissioner is hereby authorized to charge any additional filing fees required under 37 CFR 1.16 or credit any overpayment to Eastman Kodak Company Deposit Account No. 05-0225.

A duplicate copy of this sheet is enclosed. Tuell

> Attorney for Applicants Registration No. 34,278

Doreen M. Wells/sp Telephone: (716) 588-2405 Facsimile: (716) 477-1148

**ORIGINAL** 

Application Based on

Docket 81326D-W

Inventors: Jiann Hsing Chen, Joseph Anthony Pavlisko, Charles Chester Anderson and Robert Arthur Lancaster

# METHOD OF CURING A FUSER MEMBER OVERCOAT AT LOW TEMPERATURES

Commissioner for Patents, ATTN: BOX PATENT APPLICATION Washington, D. C. 20231

Express Mail Label No.: EL 485197960US

Date: June 30, 2000

25

30

5

# METHOD OF CURING A FUSER MEMBER OVERCOAT AT LOW TEMPERATURES

## CROSS REFERENCE TO RELATED APPLICATIONS

	Reference is made to commonly assigned, copending 0.5. Fatent
	Application Serial Numbers:
	by Chen et al., filed of even date herewith
	(Docket 81322), entitled "FLUOROCARBON THERMOPLASTIC RANDOM
10	COPOLYMER COMPOSITION CURABLE AT LOW TEMPERATURES".
	by Chen et al., filed of even date herewith
	(Docket 81324), entitled "METHOD OF PREPARING LOW TEMPERATURE
	CURE POLYMER COMPOSITION";
	by Chen et al., filed of even date herewith
15	(Docket 81325), entitled "FUSER MEMBER WITH LOW-TEMPERATURE-CURE
	OVERCOAT":

## FIELD OF THE INVENTION

This invention relates to a method of making fuser members useful for heat-fixing a heat-softenable toner material to a substrate. More particularly, the invention relates to a method of making materials usable as a toner release layer in a fuser member.

## BACKGROUND OF THE INVENTION

Heat-softenable toners are widely used in imaging methods such as electrostatography, wherein electrically charged toner is deposited imagewise on a dielectric or photoconductive element bearing an electrostatic latent image. Most often in such methods, the toner is then transferred to a surface of another substrate, such as, e.g., a receiver sheet comprising paper or a transparent film, where it is then fixed in place to yield the final desired toner image.

When heat-softenable toners, comprising, e.g., thermoplastic polymeric binders, are employed, the usual method of fixing the toner in place involves applying

15

20

25

30

heat to the toner once it is on the receiver sheet surface to soften it and then allowing or causing the toner to cool.

One such well-known fusing method comprises passing the toner-bearing receiver sheet through a nip formed by a pair of opposing rolls, at least one of which (usually referred to as a fuser roll) is heated and contacts the toner-bearing surface of the receiver sheet in order to heat and soften the toner. The other roll (usually referred to as a pressure roll) serves to press the receiver sheet into contact with the fuser roll. In some other fusing methods, the configuration is varied and the "fuser roll" or "pressure roll" takes the form of a flat plate or belt. The description herein, while generally directed to a generally cylindrical fuser roll in combination with a generally cylindrical pressure roll, is not limited to fusing systems having members with those configurations. For that reason, the term "fuser member" is generally used herein in place of "pressure member" in place of "pressure roll".

The fuser member usually comprises a rigid core covered with a resilient material, which will be referred to herein as a "base cushion layer." The resilient base cushion layer and the amount of pressure exerted by the pressure member serve to establish the area of contact of the fuser member with the toner-bearing surface of the receiver sheet as it passes through the nip of the fuser member and pressure members. The size of this area of contact helps to establish the length of time that any given portion of the toner image will be in contact with and heated by the fuser member. The degree of hardness (often referred to as "storage modulus") and stability thereof, of the base cushion layer are important factors in establishing and maintaining the desired area of contact.

In some previous fusing systems, it has been advantageous to vary the pressure exerted by the pressure member against the receiver sheet and fuser member. This variation in pressure can be provided, for example in a fusing system having a pressure roll and a fuser roll, by slightly modifying the shape of the pressure roll. The variance of pressure, in the form of a gradient of pressure that changes along the direction through the nip that is parallel to the axes of the rolls, can be established, for example, by continuously varying the overall diameter of the pressure roll along the direction of its axis such that the diameter is smallest at the midpoint of the axis and

15

20

25

30

largest at the ends of the axis, in order to give the pressure roll a sort of "bow tie" or "hourglass" shape. This will cause the pair of rolls to exert more pressure on the receiver sheet in the nip in the areas near the ends of the rolls than in the area about the midpoint of the rolls. This gradient of pressure helps to prevent wrinkles and cockle in the receiver sheet as it passes through the nip. Over time, however, the fuser roll begins to permanently deform to conform to the shape of the pressure roll and the gradient of pressure is reduced or lost, along with its attendant benefits. It has been found that permanent deformation (alternatively referred to as "creep") of the base cushion layer of the fuser member is the greatest contributor to this problem.

Particulate inorganic fillers have been added to base cushion layers to improve mechanical strength and thermal conductivity. High thermal conductivity is advantageous when the fuser member is heated by an internal heater, so that the heat can be efficiently and quickly transmitted toward the outer surface of the fuser member and toward the toner on the receiver sheet it is intended to contact and fuse. High thermal conductivity is not so important when the roll is intended to be heated by an external heat source.

Polyfluorocarbon elastomers, such as vinylidene fluoridehexafluoropropylene copolymers, are tough, wear resistant and flexible elastomers that have excellent high temperature resistance, but relatively high surface energies, which compromises toner release.

Fluorocarbon resins like polytetrafluoroethylene (PTFE) or fluorinated ethylenepropylene (FEP) are fluorocarbon plastics which have excellent release characteristics due to very low surface energy. Fluorocarbon resins are, however, less flexible and elastic than fluorocarbon elastomers and are therefore not suitable alone as the surface of the fuser roller.

U.S. Patent No. 4,568,275 discloses a fuser roll having a layer of fluorocarbon elastomer and a fluorinated resin powder. However, the fluorocarbon elastomer that is disclosed is water dispersible and it is known that the mixture phase separates on coating so that the fluorinated resin that is used comes to the surface of the layer.

10

15

20

25

30

U.S. Patent No. 5,253,027 discloses a fluorinated resin in a silicone elastomer. However, composites of this type exhibit unacceptable swell in the presence of silicone release oil.

U.S. Patent No. 5, 599, 631 discloses a fuser roll having a layer of a fluorocarbon elastomer and a fluorocarbon resin. The drawback of this type of material is that the fluorocarbon resin powder tends to phase separate from the fluorocarbon elastomer thereby diminishing toner release.

U.S. Patent No. 4, 853, 737 discloses a fuser roll having an outer layer comprising cured fluorocarbon elastomers containing pendant amine functional polydimethylsiloxane that are covalently bonded to the backbone of the fluorocarbon elastomer. However, the amine functional polydimethylsiloxane tends to phase senarate from the fluorocarbon elastomer.

U.S. Patent No. 5,582,917 discloses a fuser roll having a surface layer comprising a fluorocarbon-silicone polymeric composition obtained by heating a fluorocarbon elastomer with a fluorocarbon elastomer curing agent in the presence of a curable polyfunctional poly(C1-6 alkyl) siloxane polymer. However, the resulting interpenetrating network (IPN) has relatively high coefficient of friction and relatively low mechanical strength. After a period of use, the release property of the roller degrades and paper jams begin to occur.

U.S. Patent No. 5, 547,759 discloses a fuser roll having a release coating layer comprising an outermost layer of fluorocarbon resin uniquely bonded to a fluoroelastomer layer by means of a fluoropolymer containing a polyamide-imide primer layer. Although the release coating layer has relatively low surface energy and good mechanical strength, the release coating layer lacks flexibility and elastic properties and can not produce high quality of images.

In addition, sintering the fluorocarbon resin layer is usually accomplished by heating the coated fuser member to temperatures of approximately 350°C to 400°C. Such high temperatures can have a detrimental effect on the underlying base cushion layer which normally comprises a silicone rubber layer. It would be desirable to provide a fuser member with an overcoat layer comprising a fluorocarbon resin layer without depolymerizing the silicone base cushion layer on heating.

15

20

2.5

30

Polysiloxane elastomers have relatively high surface energy and relatively low mechanical strength, but are adequately flexible and elastic and can produce high quality fused images. After a period of use, however, the self release property of the roller degrades and offset begins to occur. Application of a polysiloxane fluid during roller use enhances the ability of the roller to release toner, but shortens roller life due to oil absorption. Oiled portions tend to swell and wear and degrade faster.

One type of material that has been widely employed in the past to form a resilient base cushion layer for fuser rolls is a condensation-crosslinked siloxane elastomer. Disclosure of filled condensation-cured poly(dimethylsiloxane)( "PDMS") elastomers for fuser rolls can be found, for example, in U.S. Patent. Nos. 4,373,239; 4,430,406; and 4,518,655. U.S. Patent. No. 4,970,098 to Ayala-Esquillin et al. teaches a condensation cross-linked diphenylsiloxane-dimethylsiloxane elastomer having 40 to 55 weight percent zinc oxide, 5 to 10 weight percent graphite, and 1 to 5 weight percent ceric dioxide.

A widely used siloxane elastomer is a condensation-crosslinked PDMS elastomer, which contains about 32-37 volume percent aluminum oxide filler and about 2-6 volume percent iron oxide filler, and is sold under the trade name, EC4952, by the Emerson Cummings Co., U.S.A.

It has been found that fuser rolls containing EC4952 cushion layers exhibit serious stability problems over time of use, i.e., significant degradation, creep, and changes in hardness, that greatly reduce their useful life. Nevertheless, materials such as EC4952 initially provide very suitable resilience, hardness, and thermal conductivity for fuser roll cushion layers.

U.S. Patent 5,595,823 discloses toner fusing members which have a substrate coated with a fluorocarbon random copolymer containing aluminum oxide. Although these toner fusing members have proved effective and have desirable thermal conductivity, they have a problem in that there can be toner contamination. The advantage of using the cured fluorocarbon thermoplastic random copolymer compositions is that they are effective for use with toner release agents which typically include silicone.

10

20

25

30

U.S. Patent 5,464,698 discloses toner fusing members which have a substrate coated with a fluorocarbon random copolymer containing tin oxide. Although these toner fusing members have proved effective and have desirable thermal conductivity, they have a problem in that there can be toner contamination.

Commonly-assigned U.S. Patent No. 6,041,210 describes a toner fusing member having an overcoat layer including electrically conductive fine powders having a weight percent between about 30 to 80 weight percent. Although these toner fusing members have proved effective in suppressing electrostatic charge build up, they have a problem in that there can be toner contamination.

Thus, it has been extremely difficult to provide a fuser roller with, at the same time, good wear resistance, good release property, low coefficient of friction and low oil swell when exposed to release oil. It is toward a solution to this problem that the present invention is directed. It would be desirable to provide a method for making a fuser member with an overcoat layer that includes a fluorocarbon 15 thermoplastic random copolymer that can be cured at lower temperatures while providing good mechanical properties and low toner contamination.

### SUMMARY OF THE INVENTION

The present invention provides a method of making materials for forming a toner release layer that overcome the problems described above.

The method of the present invention provides a fuser member that contains a fluorocarbon thermoplastic random copolymer having improved toner release and mechanical strength.

> The present invention is a method of making a fuser member having a support comprising the steps of:

- a) providing a support;
- b) coating from an organic solvent onto the support a coating composition comprising a fluorocarbon thermoplastic random copolymer, a curing agent having a bisphenol residue, a particulate filler containing zinc oxide, antimonydoped tin oxide particles and aminosiloxane, the fluorocarbon thermoplastic random copolymer having subunits of:

wherein

5

10

15

20

x is from 1 to 50 or 60 to 80 mole percent,

y is from 10 to 90 mole percent,

z is from 10 to 90 mole percent, and

x + y + z equal 100 mole percent.

c) curing the coating composition for 5 to 10 hours at a temperature in the range of 25-275°C.

The aminosiloxane is an amino functional polydimethyl siloxane copolymer comprising aminofunctional units selected from the group consisting of (aminoethylaminopropyl) methyl, (aminopropyl) methyl and (aminopropyl) dimethyl.

Optionally, the layer may further contain a fluorinated resin; the resin is polytetrafluoroethylene or fluoroethylene propylene and has a number average molecular weight of about 50,000 to 50,000,000. In addition, the layer may further optionally contain carbon black.

As will be demonstrated through examples, fuser members formed with a toner release layer having an unfilled fluorocarbon thermoplastic random copolymer have poor mechanical strength and toner release. However, it has been surprisingly found in the present invention that the addition of zinc oxide filler and an aminosiloxane polymer to a fluorocarbon thermoplastic random copolymer provides a fuser member having improved mechanical strength, toner release and reduced toner contamination. It was particularly surprising that the addition of antimony-doped tin oxide particles to these compositions significantly reduced the temperatures required for curing.

A further advantage of the present invention is the addition of specific release additives such as fluorinated resins to the fuser member compositions in the presence of bisphenol residue curing agent significantly improves the frictional characteristics of the fuser member.

30

25

## BRIEF DESCRIPTION OF THE DRAWINGS

10

15

20

30

FIG. 1 is a cross sectional view of a fusing member in accordance with the present invention.

### DETAILED DESCRIPTION OF PARTICULAR EMBODIMENTS

FIG. 1 shows a cross sectional view of a fuser member 10 which include fuser roller, pressure roller, oiler donor roller, oiler metering roller, preconditioning roller, etc. The support 16 is usually metallic such as stainless steel, steel, aluminum, etc.; however, the support 16 may also be made of a ceramic or plastic. The primary requisites for support 16 materials are that they provide the necessary stiffness, be able to support the force placed upon it, and be able to withstand whatever temperature to which it is subjected. Disposed above the support 16 lies one or more optional intermediate layers 14 which are characterized in the art as cushion layers. The outermost layer 12 is a toner release layer. In the event that a cushion layer 14 is not desired, then the outermost layer 12 is disposed directly over the support 16. The outermost layer 12 is the toner release layer. It includes a curing agent and a fluorocarbon random copolymer that is cured by the curing agent, the fluorocarbon random copolymer has subunits of:

$$-(CH_2CF_2)x--, -(CF_2CF(CF_3)y--, and -(CF_2CF_2)z--,$$
  
wherein

x is from 1 to 50 or 60 to 80 mole percent,

v is from 10 to 90 mole percent,

z is from 10 to 90 mole percent,

x + y + z equal 100 mole percent.

—(CH<sub>2</sub>CF<sub>2</sub>) is (vinylidene fluoride subunit ("VF<sub>2</sub>")).

5 —(CF<sub>2</sub>CF(CF<sub>3</sub>) is (hexefluoropropykene subunit ("HFP')), and

-(CF, CF,) is (tetrafluoroethylene subunit ("TFE")).

The layer further includes a bisphenol residue curing agent, a particulate filler having zinc oxide, antimony-doped tin oxide particles and aminosiloxane. The aminosiloxane is an amino functional polydimethyl siloxane copolymer comprising aminofunctional units selected from the group consisting of (aminoethylaminopropyl) methyl, (aminopropyl) methyl and (aminopropyl) dimethyl.

15

20

25

30

It is a feature of the present invention that the fuser member formed with a toner release layer that includes a zinc oxide and an aminosiloxane filled polyfluorocarbon thermoplastic random copolymer has a moderately low surface energy. By using a fluorocarbon thermoplastic polymeric composition, an improved fuser member is provided. A further advantage of the present invention is that particular optional release additives such as a fluorinated resin can be added to the fluorocarbon thermoplastic random copolymer in the presence of a bisphenol residue curing agent to improve the coefficient of friction.

In these formulas, x, y, and z are mole percentages of the individual subunits relative to a total of the three subunits (x+y+z), referred to herein as "subunit mole percentages". The curing agent can be considered to provide an additional "cure-site subunit", however, the contribution of these cure-site subunits is not considered in subunit mole percentages. In the fluorocarbon thermoplastic copolymer, x has a subunit mole percentage of from 1 to 50 or 60 to 80 mole percent, y has a subunit mole percentage of from 10 to 90 mole percent, and z has a subunit mole percentage of from 10 to 90 mole percent, and z has a subunit of the invention, subunit mole percentages are: x is from 30 to 50 or 70 to 80, y is from 10 to 20, and z is from 10 to 50; or more preferably x is from 40 to 50, y is from 10 to 15, and z is 40 to 50. In the currently preferred embodiments of the invention, x, y, and z are selected such that fluorine atoms represent at least 65 percent of the total formula weight of the VF2, HFP, and TFE subunits.

Preferably, a curable amino functional polydimethyl siloxane copolymer is used in the present invention and is cured concurrently with the fluorocarbon thermoplastic random copolymer to produce a coating suitable for use as the toner release layer of a fusing member. In accordance with the invention, coated fuser members have low energy surfaces which release toner images with minimal offset. Preferred curable amino functional polydimethyl siloxanes are bis(aminopropyl) terminated poly(dimethylsiloxane). Such oligomers are available in a series of molecular weights as disclosed, for example, by Yilgor et al., "Segmented Organosiloxane Copolymer", Polymer, 1984, V.25, pp1800-1806.

A preferred class of curable amino functional polydimethyl siloxanes, based on availability, includes those having functional groups such as aminopropyl UVICIO LO BORG

5

10

15

20

25

30

or aminoethylaminopropyl pendant from the siloxane backbone such as DMS-A11, DMS-A12, DMS-A15, DMS-A21 and DMS-A32 (sold by Gelest, Inc.) having a number- average molecular weight between 850 to 27,000. Other curable amino functional polydimethyl siloxanes which can be used are disclosed in U.S. Patent Nos. 4,853,737 and 5,157,445, the disclosures of which are hereby incorporated by reference.

Preferred compositions of the invention have a ratio of aminosiloxane polymer to fluorocarbon thermoplastic random copolymer between about 0.01 and 0.2 to 1 by weight, preferably between about 0.05 and 0.15 to 1. The composition is preferably obtained by curing a mixture comprising from about 50-80 weight percent of a fluorocarbon thermoplastic copolymer, 5-20 weight percent, most preferably about 5-10 weight percent, of a curable amino functional polydimethyl siloxane copolymer, 1-5 weight percent of a bisphenol residue, 1-20 weight percent of a zinc oxide acid acceptor type filler, 3-20 weight percent of antimony-doped tin oxide particles, 0 to 10 weight percent of carbon black, and 10-50 weight percent of a fluorinated resin release aid filler.

Curing of the fluorocarbon thermoplastic random copolymer is carried out at much lower temperatures compared to the well known conditions for curing vinylidene fluoride based fluorocarbon elastomer copolymers. For example, the cure of fluorocarbon elastomers is usually for 12-48 hours at temperatures of about 220 to 250°C. Typically, fluorocarbon elastomer coating compositions are dried until solvent free at room temperature, then gradually heated to about 230°C over 24 hours, then maintained at that temperature for 24 hours. By contrast, the cure of the fluorocarbon thermoplastic random copolymer compositions of the current invention is about 5 to 10 hours at a temperature of about 25-275°C, preferably 25-120°C, and most preferably 25-50°C.

The outer layer includes a particulate filler comprising zinc oxide. The zinc oxide particles can be obtained from a convenient commercial source, e.g., Atlantic Equipment Engineers of Bergenfield, New Jersey. In a currently preferred embodiment of the invention, the particulate zinc oxide filler has a total concentration in the outer layer of from about 1 to 20 parts per hundred parts by weight of the fluorocarbon thermoplastic random copolymer (pph). Concentrations of zinc oxide

COCCEC COCCCC

5

10

15

20

25

30

with much less than 1 part by weight may not provide the desired degree of stability to the layer. Concentrations of zinc oxide with much greater than 20 parts by weight will render the layer too stiff to provide the desired area of contact with the toner-bearing receiver sheet. In a particular embodiment of the invention, the outer layer has 3 to 10 pph of zinc oxide.

The particle size of the zinc oxide filler does not appear to be critical. Particle sizes anywhere in the range of 0.1 to 100 micrometers have been found to be acceptable. In the examples presented below the zinc oxide particles were from 1 to 40 micrometers in diameter.

The outer layers of the invention also include antimony-doped tin oxide particles. These particles can be obtained from a convenient commercial source, e.g., Keeling & Walker, Stoke-on-Trent, UK; DuPont Co; or Mitsubishi Metals Inc., Japan. In a currently preferred embodiment, the antinomy-doped tin oxide particles have a total concentration in the compositions of the invention of from about 3 to 20 parts per hundred parts by weight of the fluorocarbon thermoplastic random copolymer (pph). In a particularly preferred embodiment of the invention, the composition has 3 to 15 pph of antimony-doped tin oxide particles.

The particle size of the antimony-doped tin oxide particles does not appear to be critical. Particle sizes anywhere in the range of 0.05 to 10 micrometers are suitable. In the examples presented below the tin oxide particles were about 0.4 micrometers in diameter. The morphology or shape of the particles is not critical, for example, the particles may be essentially spherically in shape (granular) or they may be acicular in shape (e.g., a fiber or whisker).

The antimony concentration of the antimony-doped tin oxide particles is preferably 1 to 15 weight percent, most preferably 3 to 10 weight percent. In the examples presented below the antimony-doped tin oxide particles contained 6 to 9 weight percent antimony.

It was surprisingly found that the addition of the antimony-doped tin oxide particles to the fluorocarbon thermoplastic random copolymer compositions greatly reduced the temperatures needed to cure the fluoropolymer. While not wanting to be bound by theory, it is possible that the antimony acts as a catalyst for the curing reactions. Conventional fluoropolymer compositions that do not contain

10

15

20

25

30

antimony-doped tin oxide parties have curing temperatures that are typically about 220 to 280°C, while the compositions of the invention may be cured at temperatures as low as room temperature.

The compositions of the invention optionally contain a carbon black added at a concentration of 0 to 10 parts per hundred parts of the fluorocarbon thermoplastic random copolymer. Any conventional carbon black may be used, for example Thermax<sup>TM</sup> N-990 available from R.T.Vanderbilt Co.

To form the outer layer, the zinc oxide filler and antimony-doped tin oxide particles are mixed with the uncured fluorocarbon thermoplastic random copolymer, amino siloxane, a bisphenol residue curing agent, and any other optional additives, such as fluorinated resin; shaped over the base cushion, and cured. The fluorocarbon thermoplastic random copolymer is cured by crosslinking with basic nucleophile addition curing. Basic nucleophilic cure systems are well known and are discussed, for example, in U.S. Patent No. 4,272,179. One example of such a cure system combines a bisphenol residue as the curing agent and an organophosphonium salt, as an accelerator. The fluorinated resins which include polyterafluoroethylene (PTFE) or Fluoethylenepropylene (FEP) are commercially available from duPont.

The crosslinker is incorporated into the polymer as a cure-site subunit, for example, bisphenol residues. Other examples of nucleophilic addition cure systems are sold commercially as DIAK No. I (hexamethylenediamine carbamate) and DIAK No. 3 (N,N'-dicinnamylidene-1,6-hexanediamine) by duPont.

Suitable fluorocarbon thermoplastic random copolymers are available commercially. In a particular embodiment of the invention, a vinylidene fluoride-cotetrafluoroethylene co-hexafluoropropylene was used which can be represented as — (VF)(75) — (TFE) (10) - (HFP)(25) —. This material is marketed by Hoechst Company under the designation 'THV Fluoroplastics' and is referred to herein as "THV". In another embodiment of the invention, a vinylidene fluoride-cotetrafluoroethylene-co-hexafluoropropylene was used which can be represented as — (VF)(49)- (TFE) (41) — (HFP)(10) —. This material is marketed by Minnesota Mining and Manufacturing, St. Paul, Minn., under the designation "3M THV" and is

referred to herein as "THV-200A". Other suitable uncured vinylidene fluoridecohexafluoropropylenes and vinylidene fluoride-co-tetrafluoroethylenecohexafluoropropylenes are available, for example, THV-400, THV-500 and THV-300.

In general, THV Fluoroplastics are set apart from other meltprocessable fluoroplastics by a combination of high flexibility and low process temperature. With flexural modulus values between 83 Mpa and 207 Mpa, THV Fluoroplastics are the most flexible of the fluoroplastics.

The molecular weight of the uncured polymer is largely a matter of convenience, however, an excessively large or excessively small molecular weight would create problems, the nature of which are well known to those skilled in the art. In a preferred embodiment of the invention the uncured polymer has a number average molecular weight in the range of about 100,000 to 200,000.

The fuser member is constructed forming a toner release layer on an optional base cushion provided on a core comprising the steps of:

- A) providing a support;
- B) providing a mixture having:
- (i) a fluorocarbon thermoplastic random copolymer having subunits

of:

5

10

15

20

25

30

$$-(CH_2CF_2)x-$$
,  $-(CF_2CF(CF_3)y-$ , and  $-(CF_2CF_2)z-$ , wherein

x is from 1 to 50 or 60 to 80 mole percent,

y is from 10 to 90 mole percent,

z is from 10 to 90 mole percent, and

- x + y + z equals 100 mole percent
- (ii) a filler comprising zinc oxide;
- (iii) a curable amino functional polydimethyl siloxane copolymer comprising aminofunctional units selected from the group consisting of (aminoethylaminopropyl) methyl, (aminopropyl) methyl and (aminopropyl) dimethyl; and
  - (iv) antimony-doped tin oxide particles;
- (v) a bisphenol residue curing agent;
  - C) applying the mixture to the support and curing the applied mixture to crosslink the fluorocarbon thermoplastic random copolymer; and

10

1.5

20

25

30

D) curing the coating composition for 5 to 10 hours at a temperature in the range of 25°C to 275°C.

The particular conditions employed for curing within the above specified range for time and temperature may be selected based on the materials employed for the fuser member support and/or the cushion layer. For example, when the support material is a metal, higher temperatures and longer curing times may be employed. When the support is a polymer web, for example, lower curing temperatures and shorter curing times may be more appropriate. In a preferred embodiment, the substrate is either a metal or a ceramic, and the fluorocarbon thermoplastic random copolymer-containing mixtures are cured for 5 to 10 hours at a temperature of 25 to 120 °C. In another preferred embodiment, the substrate is either a plastics or a polyurethane, and the fluorocarbon thermoplastic random copolymer-containing mixtures are cured for 5 to 10 hours at a temperature of 25 to 50°C.

In cases where it is intended that the fuser member be heated by an internal heater, it is desirable that the outer layer have a relatively high thermal conductivity, so that the heat can be efficiently and quickly transmitted toward the outer surface of the fuser member that will contact the toner intended to be fused. (Depending upon relative thickness, it is generally even more desirable that the base cushion layer and any other intervening layers have a relatively high thermal conductivity. Suitable materials for the base cushion layer are discussed below.)

Some fusing systems use a release oil, such as a PDMS oil, to prevent offset, that is, to aid the roll in releasing from the toner it contacts during the fusing operation. During use, the oil is continuously coated over the surface of the fuser member in contact with the toner image. The fuser member of the invention can be used with polydimethylsiloxane, amino functionalized polydimethylsiloxane or mercapto functionalized polydimethylsiloxane release oils at normally used application rates or at reduced application rates, from about 0.5 mg/copy to 10 mg/copy (the copy is 8.5 by 11 inch 20 pound bond paper).

The outer layer of the fuser member of the invention is substantially resistant to release oil induced swelling. In a preferred embodiment of the invention, the change in size due to swelling is less than 0.1 to 1.0 percent. In an even more

preferred embodiment of the invention, the change in size due to swelling is less than 0.01 to 0.1 percent.

The thickness of the base cushion and outer layers and the composition of the base cushion layer can be chosen so that the base cushion layer can provide the desired resilience to the fuser member, and the outer layer can flex to conform to that resilience. The thickness of the base cushion and outer layers will be chosen with consideration of the requirements of the particular application intended. Usually, the outer layer would be thinner than the base cushion layer. For example, base cushion layer thicknesses in the range from 0.6 to 5.0 mm have been found to be appropriate for various applications. In some embodiments of the present invention, the base cushion layer is about 2.5 mm thick, and the outer layer is from about 25 to 30 micrometers thick.

Suitable materials for the base cushion layer include any of a wide variety of materials previously used for base cushion layers, such as the condensation cured polydimethylsiloxane marketed as EC4952 by Emerson Cummings. An example of a condensation cured silicon rubber base cushion layer is GE 4044 marketed by General Electric of Waterford, N.Y. An example of an addition cured silicone rubber is Silastic J RTV marketed by Dow Corning applied over a silane primer DC-1200 also marketed by Dow Corning.

20

25

30

5

10

15

In a particular embodiment of the invention, the base cushion is resistant to cyclic stress induced deformation and hardening. Examples of suitable materials to reduce cyclic stress induced deformation and hardening are filled condensation-crosslinked PDMS elastomers disclosed in U.S. Patent. No. 5,269,740 (copper oxide filler), U.S. Pat No. 5,292,606 (zinc oxide filler), U.S. Patent. No. 5,292,562, entitled "Fuser Roller for Fixing Toner to a Substrate", U.S. Patent No. 5,480,724, entitled "Fuser Roll for Fixing Toner to a Substrate Comprising Tin Oxide Fillers", U.S. Patent No. 5,336,539 Fuser Roll Containing Nickel Oxide Particles for Fixing Toner to a Substrate". These materials all show reasonable thermal conductivities and much less change in hardness and creep than EC4952 or the PDMS elastomer with aluminum oxide filler. Additional suitable base cushions are disclosed in U.S. Patent No. 5,466,533, entitled "Zinc Oxide Filled Diphenylsiloxane-Dimethylsiloxane Fuser Roll for Fixing Toner to a Substrate",

10

15

20

25

30

U.S. Patent No. 5,474,852, entitled "Tin Oxide Filled Diphenylsiloxane-Dimethylsiloxane Fuser Roll for Fixing Toner to a Substrate", U.S. Patent No. 5,464,703, entitled "Tin Oxide Filled Dimethylsiloxane-Fluoroalkylsiloxane Fuser Roll for Fixing Toner to a Substrate." The patents and patent applications mentioned in this paragraph are hereby incorporated herein by reference.

The support of the fuser member is usually cylindrical in shape. It comprises any rigid metal or plastic substance. Metals are preferred when the fuser member is to be internally heated, because of their generally higher thermal conductivity. Suitable core materials include, e.g., aluminum, steel, various alloys, and polymeric materials such as thermoset resins, with or without fiber reinforcement. The support which has been conversion coated and primed with metal alkoxide primer in accordance with U.S. Patent No. 5,474,821, which is hereby incorporated by reference.

The fuser member is mainly described herein in terms of embodiments in which the fuser member is a fuser roll having a support, a base cushion layer overlying the support, and an outer layer superimposed on the base cushion. The invention is not, however, limited to a roll, nor is the invention limited to a fusing member having a core bearing two layers: the base cushion layer and the outer layer. The fuser member of the invention can have a variety of outer configurations and layer arrangements known to those skilled in the art. For example, the base cushion layer could be eliminated or the outer layer described herein could be overlaid by one or more additional layers.

The invention is further illustrated by the following Examples and Comparative Example.

EXAMPLES 1 – 3

150 grams of Fluorocarbon thermoplastic random copolymer THV 200A, 1.05 grams of zinc oxide, 15.4 grams of fluorinated resin, and 4.90 grams of aminosiloxane were mixed into 230 grams of methyl ethyl ketone in a milling crock as indicated (amounts listed as parts per hundred parts of THV200A) in Table 1. THV200A is a commercially available fluorocarbon thermoplastic random copolymer which is sold by 3M Corporation. The zinc oxide particles can be obtained from

15

20

convenient commercial source, e.g., Atlantic Equipment Engineers of Bergenfield, New Jersey. The amino siloxane DMS-A21 is commercially available from Gelest, Inc. The fluorinated resin is fluoroethylenepropylene (FEP) and is commercially available from duPont. Into the above mixture antimony-doped tin oxide particles and carbon black were added and the formulations were mixed on a two-roll mill for 48 hours to form a dispersion (the amounts of the antimony-doped tin oxide particles and carbon black are given in Table 1). The antimony-doped tin oxide particles are Keeling & Walker Inc. CPM375 having an average particle size of about 0.4 □m and an antimony content of 6-9 weight %. The carbon black is Thermax™ available from R.T. Vanderbilt Co.

Each of the above dispersions were mixed with 1.05 grams (3 pph) of curative 50 (a bisphenol residue, DuPont) and roll milled for 2-3 minutes. The dispersions were then immediately cast into a film and allowed to dry for several hours. The resulting layers had a thickness of several mils. Afterwards the layers were cured using the conditions listed in Table 2.

## COMPARATIVE EXAMPLE 1

To prepare Comparative Example 1 substantially the same procedures were followed as in Example 1-3, with the following exception. As indicated in the composition listed in Table 1, Comparative Example 1 did not contain antimony-doped tin oxide or carbon black. The curing conditions employed are given in Table 2.

Table 1

Sample	THV 200A	ZnO	Amino- siloxane	FEP	CMP375 Tin oxide	Carbon black
Example 1	100	3	14	44	8	2
Example 2	100	3	14	44	8	0
Example 3	100	3	14	44	8	2
Comparative Example 1	100	3	14	44	0	0

Table 2

Sample	Post Cured	Max. Temp. For Curing
Example 1	No	25 °C
Example 2	No	25 °C
Example 3	Yes	275 °C
Comparative Example 1	No	25 °C

15

20

## DMA: Testing Method

The samples were tested on a Rheometrics RSA II Dynamic Mechanical Analyzer (DMA) and required a sample geometry of 7.5 mm X 23 mm with a thickness between 30 microns to 2000 microns. The free standing films were tested at 10Hz and a strain of 0.07%. The test was recorded over a temperature scan of -100°C to 200°C. Over the temperature scan an oscillatory strain is applied to the sample and the resulting stress is measured. These values are related to material properties by E' and E'' (Storage and Loss Moduli, respectively). As a result of DMA testing, the storage modulus (E') at three different temperatures is determined and the behavior of the material at typical toner fusing temperatures is observed.

<u>Table 3</u> Storage Modulus Versus Temperature

Sample	MPa @ 80°C	MPa @ 140°C	MPa @ 175℃
Example 1	28.0	7.05	7.05
Example 2	20.0	4.50	4.50
Example 3	11.5	4.80	4.80
Comparative Example 1	11.0	0.90	0.30

Table 3 shows a comparison between the cured fluorocarbon thermoplastic random copolymer layers of the invention and Comparative Example 1 which did not contain antimony-doped tin oxide. The comparative example, despite containing the bisphenol residue curing agent, did not cure at low temperature because it did not contain the antimony-doped tin oxide which apparently acts as an accelerator for curing. In terms of the modulus at the 175°C fusing temperature, the cured fluorocarbon thermoplastic random copolymer layers of the invention provide a significant improvement in mechanical properties at the fusing temperature. Also, the compositions of the invention can also be cured at conventional high temperatures as in Example 3 without any significant deleterious effect on properties.

The invention has been described in detail with particular reference to certain preferred embodiments thereof, but it should be appreciated that variations and modifications can be effected within the scope of the invention

#### WHAT IS CLAIMED IS:

- 1. A method of making a fuser member having a support comprising the steps of:
  - A) providing a support;
- B) coating from an organic solvent onto the support a coating composition comprising a fluorocarbon thermoplastic random copolymer, a curing agent having a bisphenol residue, a particulate filler containing zinc oxide, antimonydoped tin oxide particles and aminosiloxane, the fluorocarbon thermoplastic random copolymer having subunits of:

—(CH
$$_2$$
CF $_2$ )x—, —(CF $_2$ CF(CF $_3$ )y—, and —(CF $_2$ CF $_2$ )z—, wherein

x is from 1 to 50 or 60 to 80 mole percent,

y is from 10 to 90 mole percent,

z is from 10 to 90 mole percent,

x + y + z equals 100 mole percent; and

- C) curing the coating composition for 5 to 10 hours at a temperature in the range of 25  $^{\circ}$  C to 275  $^{\circ}$  C.
- The method of claim 1 wherein the aminosiloxane is an amino functional polydimethyl siloxane copolymer.
- 3. The method of claim 2 wherein the amino functional polydimethyl siloxane copolymer comprises amino functional units selected from the group consisting of (aminoethylaminopropyl) methyl, (aminopropyl) methyl and (aminopropyl) dimethyl.
- 4. The method of claim 1 wherein the amino siloxane has a total concentration in the layer of from 1 to 20 parts by weight per 100 parts of the fluorocarbon thermoplastic random copolymer.

- 5. The method of claim 1 wherein the aminosiloxane has a total concentration in the layer of from 5 to 15 parts by weight per 100 parts of the fluorocarbon thermoplastic random copolymer.
- 6. The method of claim 1 wherein the aminosiloxane has a total concentration in the layer of from 10 to 15 parts by weight per 100 parts of the fluorocarbon thermoplastic random copolymer.
- 7. The method of claim 1 wherein the zinc oxide has a total concentration in the layer of from 1 to 20 parts by weight per 100 parts of the fluorocarbon thermoplastic random copolymer.
- 8. The method of claim 1 wherein the zinc oxide has a total concentration in the layer of from 3 to 15 parts by weight per 100 parts of the fluorocarbon thermoplastic random copolymer.
- The method of claim 2 wherein the fluorocarbon thermoplastic random copolymer is cured by bisphenol residues.
- $10.\,$  The method of claim 1 further including a cushion layer between the core and the layer.
- 11. The method of claim 1 wherein the fluorocarbon thermoplastic random copolymer is nucleophilic addition cured.
- 12. The method of claim 1 wherein x is from 30 to 50 mole percent, y is from 10 to 90 mole percent, and z is from 10 to 90 mole percent.
- 13. The method of claim 1 wherein x is from 40 to 50 mole percent and y is from 10 to 15 mole percent.
  - 14. The method of claim 1 wherein z is greater than 40 mole percent.

- 15. The method of claim 1 wherein the antimony-doped tin oxide particles have a total concentration of from 3 to 20 parts by weight per 100 parts of the fluorocarbon thermoplastic random copolymer.
- 16. The method of claim 1 wherein the antimony-doped tin oxide particles comprise 3 to 10 weight percent antimony.
- 17. The method of claim 1 wherein the fluorocarbon thermoplastic random copolymer further comprises a fluorinated resin.
- 18. The method of claim 17 wherein the fluorinated resin has a number average molecular weight of between 50,000 to 50,000,000.
- 19. The method of claim 17 wherein the ratio of fluorocarbon thermoplastic random copolymer to fluorinated resin is between 1:1 to 50:1.
- 20. The method of claim 17 wherein the fluorinated resin is polytetrafluoroethylene or fluoroethylenepropylene.

15

wherein

## ABSTRACT OF THE DISCLOSURE

A method of making a fuser member having a support comprising the steps of:

- A) providing a support;
  - B) coating from an organic solvent onto the support a coating composition comprising a fluorocarbon thermoplastic random copolymer, a curing agent having a bisphenol residue, a particulate filler containing zinc oxide, an aminosiloxane, and antimony-doped tin oxide particles, the fluorocarbon thermoplastic random copolymer having subunits of:
  - —( $CH_2CF_2$ )x—, —( $CF_2CF(CF_3)$ y—, and —( $CF_2CF_2$ )z—,

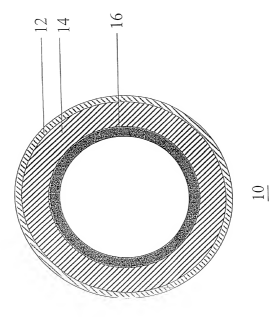
x is from 1 to 50 or 60 to 80 mole percent,

y is from 10 to 90 mole percent,

z is from 10 to 90 mole percent,

x + y + z equals 100 mole percent; and

C) curing the coating composition for 5 to 10 hours at a temperature in the range of 25°C to 275°C.



7ig. 1

combined Declaration F	or Patent Ap	plication and	Power of Attorney			81326D-		
As below named invento	r, l hereby declare	that:						
ty residence, post office address at	nd citizenship are a	s stated below next	to my name,					
believe I am the original, first an	d sole inventor (if	only one name is l	isted below) or an original,	first and joir	t invent	or (if plural r	ames are	listed
elow) of the subject matter which	is claimed and for	which a patent is sou	ight on the invention entitle	1:				
METHOD OF CURIN	G A FUSEI	R MEMBER	OVERCOAT AT	LOW T	EMPI	ERATUI	RES	
he specification of which (check of	only one item below	w):						
is attached hereto.								
was filed as United States		al No. on and						
was amended on (if applic								
was filed as PCT internation	onal application N	lumber on and w	as amended under PCT A	rticle 19 on	(if applie	cable).		
hereby state that I have reviewed :	and understand the	contents of the abo	ve-identified specification, i	ncluding the	laims, a	amended by	any ame	ndmen
eferred to above								
acknowledge the duty to disclose	to the U.S. Patent	& Trademark Offic	e all information known to	ne to be mate	rial to pa	tentability as	defined	in Title
17 Code of Federal Regulations, §	1.56.							
berely, claim foreign priority here	efits under Title 35	i, United States Cod	e, §119 of any foreign appli	cation(s) for p	atent or	inventor's cer	tificate or	r of an
CT international application(s) de	signating at least of	one country other th	an the United States of Ame	rica listed bel	ow and l	ave also iden	tified bel	ow an
breign applications(s) for patent of	or inventor's certifi	cate or any PCT in	ternational application(s) de	signating a lo	ast one	country other	than the	Unite
States of America filed by me on th	ne same subject ma	tter having a filing	late before that of the applic	ation(s) of wh	ich prio	ity is claimed	l:	
PRIOR FOREIGN/PCT APPLIC	ATION(S) AND	ANY PRIORITY C	LAIMS UNDER 35 U.S.C.	119:				
COUNTRY (V PCT, anthode PCT)	APPLI	CATION NUMBER	DATE OF FILING (Say month year)			TROPITY CLAIMED UNI	ER SS USC 511	MO
irror, assault or						YES		
						YES		NO
			1				1	
					n(s) liste	ves d below:		NO
I hereby claim the benefit under Ti	CATION(S) AND							NO
	CATION(S) AND			. §119 (e):				NO
PRIOR PROVISIONAL APPLIC	CATION(S) AND			. §119 (e):				NO
PRIOR PROVISIONAL APPLIC	CATION(S) AND	ANY PRIORITY C	LAIMS UNDER 35 U.S.C	. §119 (е):	ATE	d below:		
PRIOR PROVISIONAL APPLIC	CATION(S) AND	States Code, \$120	LAIMS UNDER 35 U.S.C	s application	a(s) or F	d below:  CT internation this application	on is not	ication
PRIOR PROVISIONAL APPLIC	r Title 35, United	ANY PRIORITY C	LAIMS UNDER 35 U.S.C  of any prior United State of a state of the stat	es application	u(s) or F laims of the duty	d below:  CT internative this application to disclose to	on is not to the U.S.	icatior disclor
PRIOR PROVISIONAL APPLIC PROVISIONAL APPLIC PROVISIONAL APPLIC PROVISIONAL APPLICATION APP	Title 35, United merica that is/are in the manner prov	ANY PRIORITY C	LAIMS UNDER 35 U.S.C  of any prior United State of are as the subject matter of agraph of Title 35, §112, 1 ability as defined in Title 3	es application each of the cucknowledge 7, Code of Fe	(s) or F laims of the duty	d below:  CT internation this application of isolose to gulations §1	on is not to the U.S.	icatior disclor
PRIOR PROVISIONAL APPLIC PROMISES OF THE PROVISIONAL OF THE PROVISION OF T	r Title 35, United the transport of the prior application.	States Code, \$120 listed below and, insided by the first pare material to patent ion(s) and the nation	LAIMS UNDER 35 U.S.C  of any prior United State to far as the subject matter of agraph of Title 35, \$112, 1 ability as defined in Title 3 and or PCT international filtred.	es application feach of the cacknowledge 7, Code of Feg date of this	u(s) or F laims of the duty deral Re applicat	d below:  CT internation this application disclose to gulations §1 ion:	on is not the U.S.	icatior disclor
I hereby claim the benefit under designating the United States of A in that/those prior applications(s). 'Tademark Office all information available between the filing date or PRIOR US APPLICATIONS O	r Title 35, United the transport of the prior application.	States Code, \$120 listed below and, insided by the first pare material to patent ion(s) and the nation	LAIMS UNDER 35 U.S.C  of any prior United State to far as the subject matter of agraph of Title 35, \$112, 1 ability as defined in Title 3 and or PCT international filtred.	es application feach of the cacknowledge 7, Code of Feg date of this	u(s) or F laims of the duty deral Re applicat	d below:  CT internation this application disclose to gulations §1 ion:	on is not the U.S.	icatior disclor
PRIOR PROVISIONAL APPLIC PROMISES OF THE PROVISIONAL OF THE PROVISION OF T	Title 35, United in the price and the price	States Code, \$12( listed below and, instided by the first parameter of the state of	LAIMS UNDER 35 U.S.C  of any prior United State to far as the subject matter of agraph of Title 35, \$112, 1 ability as defined in Title 3 and or PCT international filtred.	es application feach of the cacknowledge 7, Code of Feg date of this	u(s) or F laims of the duty deral Re applicat	CT internative this application of sicoses to guarantee to gualations \$1 ion:	on is not the U.S. .56, whic	icatior disclor
PRIOR PROVISIONAL APPLICATIONS OF PRIOR PROVISIONAL APPLICATION APPLICATION OF PRIOR	Title 35, United merica that is/are in the manner proving when the manner proving the properties of the prior application	ANY PRIORITY C  States Code, \$125 isted below and, intrided by the first pare material to patent ionn(s) and the micn(s) and the notion(s)	LAIMS UNDER 35 U.S.C  of any prior United State to far as the subject matter of agraph of Title 35, \$112, 1 ability as defined in Title 3 and or PCT international filtred.	es application feach of the cacknowledge 7, Code of Feg date of this	(s) or F laims of the duty ederal Re applicat R BENE	d below:  CT internation this application disclose to gulations §1 ion:	on is not the U.S. .56, which	icatior disclor Paten h beca
I hereby claim the benefit under designating the United States of A in that/those prior applications(s). 'Tademark Office all information available between the filing date or PRIOR US APPLICATIONS O	Title 35, United merica that is/are in the manner proving when the manner proving the properties of the prior application	ANY PRIORITY C  States Code, \$125 isted below and, intrided by the first pare material to patent ionn(s) and the micn(s) and the notion(s)	LAIMS UNDER 35 U.S.C.  Of any prior United State tofar as the subject matter of agraph of Title 5, \$112, 1, 1 ability as defined in Title 3 nal or PCT international filli ATIONS DESIGNATING 1	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	icatior disclor Paten h beca
PRIOR PROVISIONAL APPLICATIONS OF PRIOR PROVISIONAL APPLICATION APPLICATION OF PRIOR	Title 35, United merica that is/are in the manner proving when the manner proving the properties of the prior application	ANY PRIORITY C  States Code, \$125 isted below and, intrided by the first pare material to patent ion(s) and the motion(s) and the motion state of the	LAIMS UNDER 35 U.S.C.  Of any prior United State tofar as the subject matter of agraph of Title 5, \$112, 1, 1 ability as defined in Title 3 nal or PCT international filli ATIONS DESIGNATING 1	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	icatior disclor Paten h beca
PRIOR PROVISIONAL APPLICATIONS OF PRIOR PROVISIONAL APPLICATION APPLICATION OF PRIOR	Title 35, United merica that is/are in the manner proving when the manner proving the properties of the prior application	ANY PRIORITY C  States Code, \$125 isted below and, intrided by the first pare material to patent ion(s) and the motion(s) and the motion state of the	LAIMS UNDER 35 U.S.C.  Of any prior United State tofar as the subject matter of agraph of Title 5, \$112, 1, 1 ability as defined in Title 3 nal or PCT international filli ATIONS DESIGNATING 1	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
PRIOR PROVISIONAL APPLIC PROVISIONAL APPLICATION APPLICATION APPLICATION NUMBER  U.S. APPLICATIO	Title 35, United merical that is a control to the c	States Code, \$12(sixed below and, in indiced by the first of the control of the c	LAIMS UNDER 35 U.S.C.  Of any prior United State tofar as the subject matter of agraph of Title 5, \$112, 1, 1 ability as defined in Title 3 nal or PCT international filli ATIONS DESIGNATING 1	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
PRIOR PROVISIONAL APPLIC PROVISIONAL APPLICATION APPLICATION APPLICATION NUMBER  U.S. APPLICATIO	Title 35, United merica that is/are in the manner proving when the manner proving the properties of the prior application	States Code, \$12(sixed below and, in indiced by the first of the control of the c	LAIMS UNDER 35 U.S.C.  Of any prior United State tofar as the subject matter of agraph of Title 5, \$112, 1, 1 ability as defined in Title 3 nal or PCT international filli ATIONS DESIGNATING 1	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
Il hereby claim the benefit unde designating the United States of A in that/hose prior applications(s). The designating the United States of A in that/hose prior applications(s). Thademark Office all information available between the films date of PRIOR US APPLICATIONS Of 36USC\$120:  U.S. APPLICATION NUMBER OF A INC. APPLICATION NUMB	TAPPLICATIONS DES	ANY PRIORITY C  States Code, \$12( listed below and, in- rided by the first pare material to patent cion(s) and the natio  TIONAL APPLICA  ATIONS  U.S.  SIGNATING THE U.S.	LAIMS UNDER 35 U.S.C.  Of any prior United State of the subject matter of ungraph of Title 35, \$112, 1: ability as defined in Title 3 and or PCT international filling and or PCT international filling the subject of t	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
PRIOR PROVISIONAL APPLIC PROVISIONAL APPLICATION APPLICATION APPLICATION NUMBER  U.S. APPLICATIO	Title 35, United merical that is a control to the c	ANY PRIORITY C  States Code, \$12( listed below and, in- rided by the first pare material to patent cion(s) and the natio  TIONAL APPLICA  ATIONS  U.S.  SIGNATING THE U.S.	LAIMS UNDER 35 U.S.C.  D of any prior United State of the subject matter of anymph of Title 55, \$112, 11. abdility as defined in Title 3 and or PCT international filling the subject of t	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
Il hereby claim the benefit unde designating the United States of A in that/hose prior applications(s). The designating the United States of A in that/hose prior applications(s). Thademark Office all information available between the films date of PRIOR US APPLICATIONS Of 36USC\$120:  U.S. APPLICATION NUMBER OF A INC. APPLICATION NUMB	TAPPLICATIONS DES	ANY PRIORITY C  States Code, \$12( listed below and, in- rided by the first pare material to patent cion(s) and the natio  TIONAL APPLICA  ATIONS  U.S.  SIGNATING THE U.S.	LAIMS UNDER 35 U.S.C.  Of any prior United State of the subject matter of ungraph of Title 35, \$112, 1: ability as defined in Title 3 and or PCT international filling and or PCT international filling the subject of t	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos
Il hereby claim the benefit unde designating the United States of A in that/hose prior applications(s). The designating the United States of A in that/hose prior applications(s). Thademark Office all information available between the films date of PRIOR US APPLICATIONS Of 36USC\$120:  U.S. APPLICATION NUMBER OF A INC. APPLICATION NUMB	TAPPLICATIONS DES	ANY PRIORITY C  States Code, \$12( listed below and, in- rided by the first pare material to patent cion(s) and the natio  TIONAL APPLICA  ATIONS  U.S.  SIGNATING THE U.S.	LAIMS UNDER 35 U.S.C.  Of any prior United State of the subject matter of ungraph of Title 35, \$112, 1: ability as defined in Title 3 and or PCT international filling and or PCT international filling the subject of t	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
Il hereby claim the benefit unde designating the United States of A in that/hose prior applications(s). The designating the United States of A in that/hose prior applications(s). Thademark Office all information available between the films date of PRIOR US APPLICATIONS Of 36USC\$120:  U.S. APPLICATION NUMBER OF A INC. APPLICATION NUMB	TAPPLICATIONS DES	ANY PRIORITY C  States Code, \$12( listed below and, in- rided by the first pare material to patent cion(s) and the natio  TIONAL APPLICA  ATIONS  U.S.  SIGNATING THE U.S.	LAIMS UNDER 35 U.S.C.  Of any prior United State of the subject matter of ungraph of Title 35, \$112, 1: ability as defined in Title 3 and or PCT international filling and or PCT international filling the subject of t	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca
I hereby claim the benefit unde designating the United States of A in the Market of the Control	TAPPLICATIONS DES	ANY PRIORITY C  States Code, \$12( listed below and, in- rided by the first pare material to patent cion(s) and the natio  TIONAL APPLICA  ATIONS  U.S.  SIGNATING THE U.S.	LAIMS UNDER 35 U.S.C.  Of any prior United State of the subject matter of ungraph of Title 35, \$112, 1: ability as defined in Title 3 and or PCT international filling and or PCT international filling the subject of t	es application each of the cacknowledge 7, Code of Frag date of this	(s) or F laims of the duty ederal Re applicat R BENE	CT internation this application of sicological to disclose to disclose to the control of the con	on is not the U.S. .56, which	ication disclos Paten h beca

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith (List name and registration number)

Lawrence P. Kessler, Registration No. 24,637 Norman Rushefsky, Registration No. 25,606 Doreen M. Wells, Registration No. 34,278 Sarah Meeks Roberts, Registration No. 33,447

Sei	nd Correspo	Lawrence P.		Direct Telephone Calls to: (name and telephone number) Lawrence P. Kessler		
		NexPress So	(716) 253-0123			
		1447 St. Pau	l Street			
			New York 14653-7001	FAX: (716) 726-9178		
Т	FULL NAME OF	FAMILY NAME	FIRST GIVEN NAME	SECOND GIVEN NAME		
2	INVENTOR	Chen	Jiann	Н.		
٦	RESIDENCE & CITIZENSHIP	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP USA		
" L	CITIZENSAP	Fairport	New York 14450 USA	STATE & ZIP CODE (COUNTRY)		
٠ [	BUSINESS ADDRESS	BUSINESS ADDRESS	1447 St. Paul Street, Rochester	New York 14653-7001 USA		
	ADDITION	NexPress Solutions LLC	FIRST GIVEN NAME	SECOND GIVEN NAME		
2	FULL NAME OF INVENTOR	FAMILY NAME	Joseph	A.		
~ [		Pavlisko	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP		
۰۱	RESIDENCE & CITIZENSHIP	CITY Pittsford	New York 14534 USA	USA		
1		PITISTOTO BUSINESS ADDRESS	CITY	STATE & ZIP CODE (COUNTRY)		
2	ADDRESS	NexPress Solutions LLC	1447 St. Paul Street, Rochester	New York 14653-7001 USA		
-		FAMILY NAME	FIRST GIVEN NAME	SECOND GIVEN NAME		
2	PULL NAME OF INVENTOR	Anderson	Charles	C.		
ı		CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP		
0	RESIDENCE A CITIZENSHIP	Penfield	New York 14526 USA	USA		
-	BUSINESS	BUSINESS ADDRESS	CITY	STATE & ZIP CODE (COUNTRY)		
3	ADDRESS	Eastman Kodak Company	343 State Street, Rochester	New York 14650 USA		
$\dashv$	FLU NUME OF	FAMILY NAME	FIRST GIVEN NAME	SECOND GIVEN NAME		
2	INVENTOR	Lancaster	Robert	A.		
- 1	RESIDENCE &	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP		
٥	CITIZENSHIP	Hilton	New York 14468 USA	USA		
.	BUSINESS	BUSINESS ADDRESS	CITY	STATE & ZIP CODE (COUNTRY)		
4	ADDRESS	Eastman Kodak Company	343 State Street, Rochester	New York 14650 USA		
2	FULL NUMBERS	FAMILY NAME	FIRST GIVEN NAME	SECOND GIVEN NAME		
-	RESIDENCE &	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP		
0	RESIDENCE & CITIZENSHIP					
5	BUSINESS ADDRESS	BUSINESS ADDRESS	CITY	STATE & ZIP CODE (COUNTRY)		
2	FULL NIME OF INVENTOR	FAMILY NAME	FIRST GIVEN NAME	SECOND GIVEN NAME		
	RESIDENCE &	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP		
6	BUSINESS	BUSINESS ADDRESS	CITY	STATE & ZIP CODE (COUNTRY)		

hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and behef are beheved to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine of imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may joopardize the validity of the application or any patent issuing thereon.

	SIGNATURE OF INVENTOR 282	II SIGNATURE OF INVENTOR 203 .
SIGNATURE OF INVENTOR 201	SIGNATURE OF INVENTOR 202	SIGNATURE OF INVENTOR 203
Trisis Chen		Chy othertre
1 2000	DATE	DATE
DATE		54.15
June 29. 2000	Gless 29 word	Sune 29 2000
SIGNATURE OF INVENTOR 204	SIGNATURE OF INVENTOR 205	SIGNATURE OF INVENTOR 206
Folit Ci. Lancati		
DATE	DATE	DATE
June 29th 2000		